

US EPA ARCHIVE DOCUMENT



# Screening Chemicals in Commerce to Identify Possible Persistent and Bioaccumulative Chemicals: New Results and Future Work

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## Outline



- The “existing chemical” universe
- Assessments/categorization of chemicals in commerce
- Selected chemicals with P, B and POP characteristics from a combined USEPA/Canadian DSL dataset
- Review of Low/Medium Production Chemicals and chemicals that are potential biomagnifiers in air-breathing organisms
- Toxicity Reviews
- Future Pollution Prevention Studies
- Future challenges



# The Universe of Chemicals



## *Globally*

- With CAS numbers: **33,760,000** organic + inorganic substances (Feb/08)
- Commercially available: **19,184,000**
- Inventoried/regulated chemicals: **246,000**
- In commerce in USA, EU, Japan: **~100,000 with 30,000 > 1 t/yr**
- **Tracked by US EPA's Toxics Release Inventory: 650**
- **Routinely measured in environmental media: <1000**

## *In Canada*

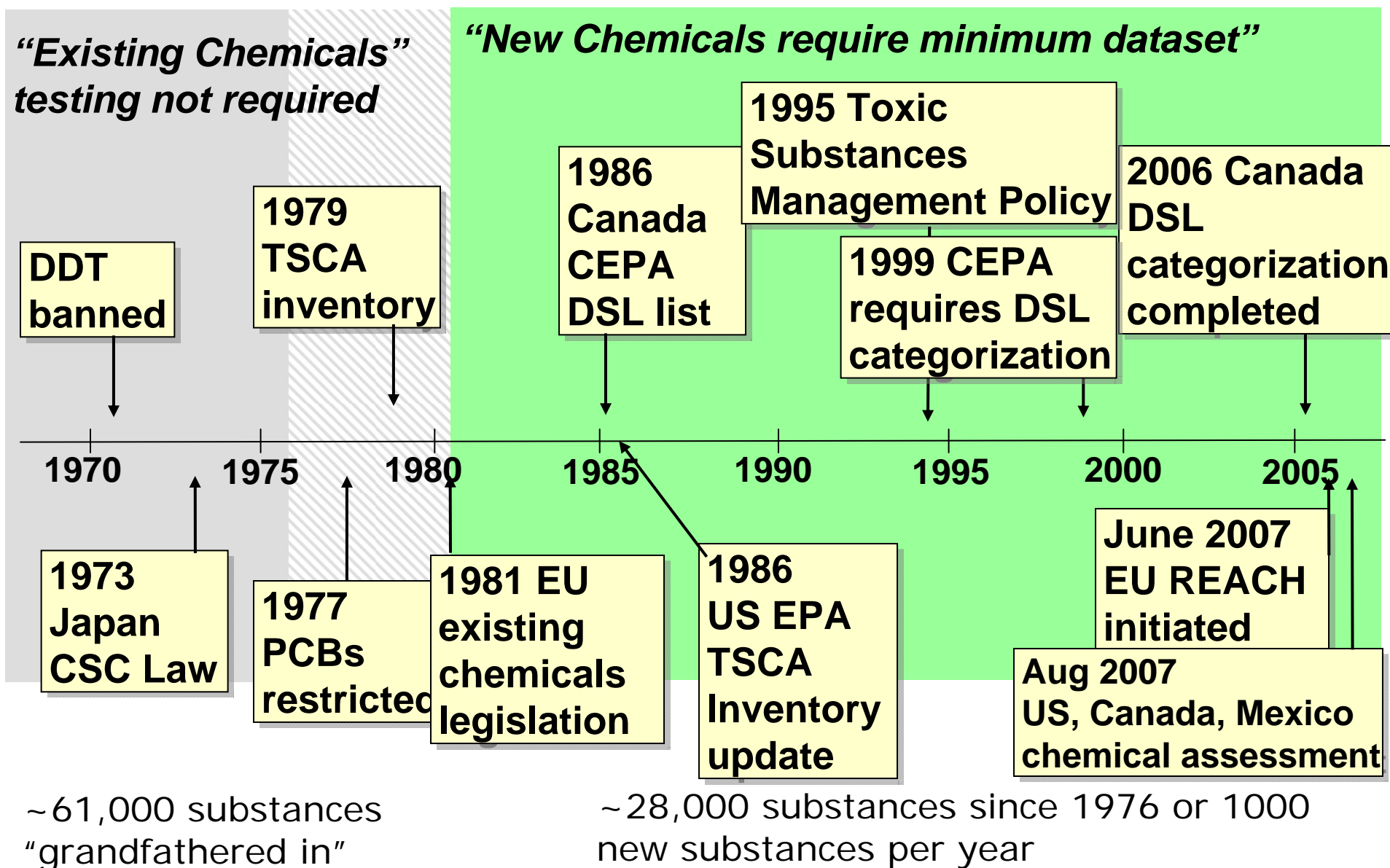
- Domestic Substances List (DSL): 24,700 (established in 1986)
- DSL & Non DSL: 70,000 substances (as of 2006)

## *In USA*

- Toxic Substances Control Act (TSCA) and TSCA Inventory update: originally 62,000 (1976), now ~82,000 substances

US FDA 1906	FIFRA (1947) 1970 pesticides
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## Regulating chemicals in commerce





## Screening of *existing chemical* lists is widening the data available for identifying potential P,B & T substances



- **TSCA Inventory Screening (US EPA)**
  - Inventory Update Rule (IUR) lists chemicals >454 t (10,000 lbs)/yr
  - Updates 1986, 1990, 1994, 1998, 2002, 2006
  - 13,750 substances (all years)
- **US EPA High Production Volume Challenge Program**
  - 2800 chemicals
  - >1,000,000 lbs/yr
- **European "REACH" (Registration, Evaluation & Authorization)**
  - Evaluation of ~5000 substances with production >100 t/yr
  - Registration of ~30,000 substances produced at >1 t/yr
  - Came into force June 2007



## Environment Canada's Domestic Substances List Categorization



- Mandated under CEPA 1999; completed September 2006
- 23,000 substances in production/use >100 kg/yr in 1986
- Screening of 11,300 organics with QSAR predicted properties + 1436 organics of unknown composition (UVCBs)
- Screening criteria: Persistent (P) or bioaccumulative (B), in accordance with Canadian P and B guidelines, and inherently toxic (iT) to humans or to non-human organisms, as determined by lab or other studies
- 4300 chemicals categorized for further assessment under the Chemicals Management Plan (CMP)



## Goals of Our Study



- Develop a North American rather than Canadian list of potentially PB&T chemicals
  - Greater relevance to the Great Lakes and trans-boundary long range transport than CMP priorities
- Using Quantitative Structure-Property relationships, and scientific judgment, identify chemicals in commerce that may be P and B and have not been previously measured in environmental media
- Assess whether selected chemicals can be analyzed by existing methods in use for POPs and new PB&T chemicals in the Great Lakes and the Arctic
- Analyze use and potential environmental release of new emerging contaminants
- Look for pollution prevention opportunities



## Development of a Combined Canadian and US database of chemicals in commerce (Howard and Meyland 2007)



Source	No. substances	Reporting threshold	Reporting date
US EPA High production volume (HPV) program*	3549	1,000,000 lbs/yr (454 t/yr)	Post-1990
US EPA TSCA Inventory update rule (IUR) web site**	14,458 organics (combined HPV and EHPVs)	>10,000 lbs/yr (4540 kg/yr)	IUR reporting years; 1986 to 2002
Canadian DSL categorization***	11,317 organics	>100 kg	Mid-1980s
UVCBs**** (1400 on the DSL)	3059 organics	>100 kg	Mid-1980s
<b>Total (after duplicates removed)</b>	<b>22,043</b>		

\*available from <http://www.epa.gov/HPV/hpvchmlt.htm>

\*\* available from <http://www.epa.gov/oppt/iur>

\*\*\* available from Environment Canada - <http://www.ec.gc.ca/substances/>

\*\*\*\* UVCB = Unknown, of Variable Composition, or of Biological Origin – organic chemicals



# Persistence and Bioaccumulation Characteristics of the 22,043 Chemicals Estimated Using EPI Suite Version 3.12



Characteristics*	No.	%	Notes
$\log K_{ow} > 5$	4239	19%	Indicates tendency to adsorb to sediments and to bioaccumulate
BCF > 2000	924	4.6%	Bioaccumulation from water exposure – does not include biomagnification
BCF > 5000	566	2.8%	
BCF > 50,000	19	0.1%	
AO* half-life > 2 days	1973	10%	AO half-life indicates stability to atmospheric oxidation and potential long range transport
AO half-life > 10 days	840	4%	
$\log K_{aw} > -5$ <u>and</u> $\log K_{aw} < -1$	6515	32%	$K_{aw}$ describes air-water partitioning. Compounds with $\log K_{aw} > -5$ & $< -1$ are "hoppers"
$\log K_{ow} \sim 2-5$ <u>and</u> high $\log K_{oa}$ $\sim 6-12$	2000	10%	Biomagnification in air-breathing organisms (Kelley et al. 2007)

\* $K_{ow}$  = octanol water partition coefficient

BCF = bioconcentration factor predicted with EPI suite software

AO = atmospheric oxidation half-life

$K_{aw}$  = air-water partition coefficient



## Further Prioritization Based on Lessons Learned from POPs in the Great Lakes and in the Arctic



1. High bioaccumulation/biomagnification potential, i.e., in top predators
2. Persistence – sequestered in bottom sediments in the open lakes implying a low rate of biodegradation
3. Long range transport potential i.e., found in mid-lake, in Lake Superior and remote lakes such as Siskiwit Lake
4. Quantity in use and potential for emissions i.e., open use or as an additive vs. as a chemical intermediate

Selection Characteristics	#	Notes
Predicted BCF > 1000, Atmospheric Oxidation > 1 day, and Log K <sub>ow</sub> > -5 and < -1	105	Using EPIsuite. Mainly chemicals with LRT potential
By chemical class (Br, Cl, F, I, Si, cyclic HCs) and considering biodegradability	324	By expert judgment – includes chemicals and their degradation products with low LRT but potential for persisting in sediments and in the water column
Total	429	<b>70% halogenated; 10% siloxanes</b>



## Information on Measurement and Analyzability of the 429 Substances



<b>Analysable</b>	<b>Well monitored in the GL region and Arctic (i.e., programs such as IADN, NCP)</b>	<b>Chemicals that may have been analysed in any GL &amp; Arctic measurement studies</b>	<b>Analyzable using existing methods for neutral POPs or other neutrals such as pesticides</b>	<b>Analyzable by LC-MS/MS ESI mode (anionic) or positive CI mode</b>
Yes	16	83	280	46
% Yes	4%	19%	65%	11%
No	413	346	116	
Maybe			33	11

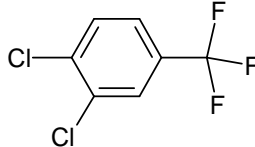
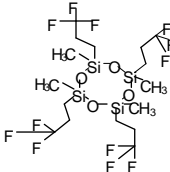
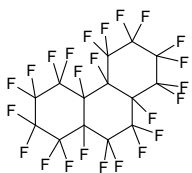
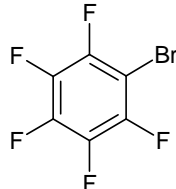
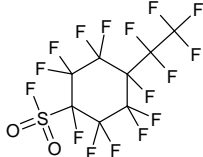


## Further Prioritization of Low-Medium Production Volume and Potential Biomagnifiers in Air-Breathing Organisms

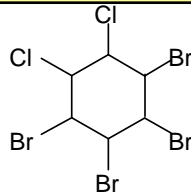
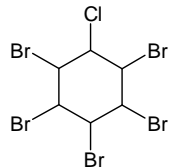
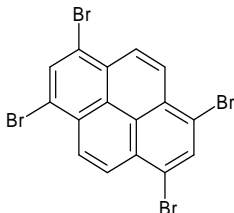
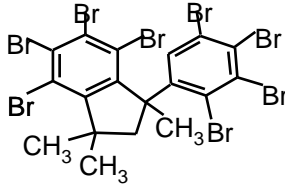
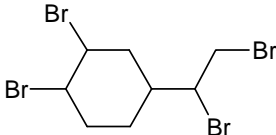


- Low-Medium Production Volume
  - Low: 10,000-500,000 lbs/yr
  - Medium: 500,000-1,000,000 lbs/yr
  - Total of low and medium = 9378 chemicals
    - » Scientific judgment screening = 270 chemicals
- Biomagnifiers in Air-Breathing Organisms
  - Low  $K_{ow} = \sim 2-5$  and  $\text{Log } K_{oa} \ 6-12 = 2000$  chemicals
    - » Scientific judgment screening = 95 chemicals
- Total Chemicals (429 + 184) - 613
  - 62% are halogenated,
  - 27% are hydrocarbons or N or S containing HCs
  - 8% are siloxanes
  - 3% are P containing.

## Some Semi-Volatile, Medium or High Production, Chlorinated or Fluorinated Organics of Potential Interest Due to (1) Predicted Persistence and/or Bioaccumulation and (2) Ease of Analysis

Substances	Predicted properties	Prod'n/yr (USA, lbs)	Structure	Notes
<b>1,2-dichloro-4-(trifluoromethyl)-benzene</b>	$\text{Log } K_{ow} = 4.2$ $\text{AOT}_{1/2} > 100 \text{ d}$	1-10 M (2002)		Very P, highly volatile. Intermediate in pesticide production
<b>Trifluoropropyl methyl cyclotetrasiloxane</b>	$\text{Log } K_{ow} = 10.7$ $\text{AOT}_{1/2} > ?$	0.5-1.0 M (2002)		P, low predicted BCF
<b>Perfluoroperhydrophenanthrene</b>	$\text{Log } K_{ow} = 9.6$ $\text{AOT}_{1/2} = ?$	0.010-0.5 M (2002)		Looks very P and B
<b>Tetrafluorobromobenzene</b>	$\text{Log } K_{ow} = 3.9$ $\text{AOT}_{1/2} = 85 \text{ d}$	0.010-0.5 M (1998)		Looks P and B
<b>Cyclohexane-sulfonyl fluoride and sulfonate</b>	$\text{Log } K_{ow} = 5.9$ $\text{AOT}_{1/2} = 77 \text{ d}$	0.010-0.5 M (2002)		P and possibly B as sulfonate deg'n product

## Some Brominated Organics of Potential Interest Due to Production Volume, High Predicted P & B and Ease of Analysis

Substances	Predicted properties	Prod'n/yr (USA, lbs and last year rep'd	Structure
<b>Tetrabromo-dichloro-cyclohexane</b>	$\text{Log } K_{\text{ow}} = 4.6$ P & B	>10-500 k (1998)	
<b>Pentabromo-6-chloro-cyclohexane</b>	$\text{Log } K_{\text{ow}} = 4.7$ P & B	>10-500 K (2002)	
<b>1,3,6,8- tetrabromopyrene</b>	$\text{Log } K_{\text{ow}} = 8.5$ Persistent May not be B	>0.5-1M (2002)	
<b>Octabromo-1,1,3-trimethyl-3-phenyl indan</b>	$\text{Log } K_{\text{ow}} \sim 10$ P & B	>0.5-1 M (2002)	
<b>1,2-dibromo-4-(1,2-dibromoethyl)cyclohexane</b>	$\text{Log } K_{\text{ow}} = 5.2$ P & B	>10-500 K (2002)	



## Toxicity Estimates of Priority Chemicals



- 429 chemicals in commerce previously identified as having potential for persistence and bioaccumulation in the Great Lakes
- Further evaluated to identify and estimate whether these compounds are toxic to aquatic organisms and to mammals utilizing:
  - Analog Identification Methodology (AIM)
  - ECOSAR
  - OncoLogic



# AIM



- EPA is currently developing the AIM tool to identify close analogs that have measured data
- Designed to help identify publicly available, experimental toxicity data on closely related chemical structures
- AIM database contains 31,031 potential analogs with publicly available toxicity data
- Experimental data sources Indexed
  - On-Line Databases
    - » TSCATS, HSDB, IRIS
  - U.S. Government Documents
    - » NTP, ATSDR, HPV Challenge Program
  - Other Sources
    - » DSSTox, RTECS, IUCLID, AEGLS



## AIM Results



- The AIM tool was run to identify if a chemical is in one of the approximately 45 EPA Chemical Classes used by the New Chemicals Program under TSCA
  - These 45 chemical classes have been identified as being of potential concern for human health effects
- 277 chemicals were included in the 45 chemical classes
- 152 chemicals were not in the 45 chemical classes



## ECOSAR



- ECOSAR is a computerized program for aquatic toxicity estimates that is currently used by EPA's Office of Pollution Prevention and Toxics (OPPT)
- Part of the EPISuite™ software — provides estimates of potential for aquatic toxicity based up  $K_{ow}$  and chemical class
- To date, over 150 SARs have been developed for more than 50 chemical classes
- This analysis involves the application of SARs (Structure Activity Relationships) to predict the aquatic toxicity of chemicals ( $LC_{50}$ ,  $EC_{50}$ , chronic, etc.) for various aquatic organisms (fish, daphnid, algae, etc.)



## ECOSAR Results



- The most toxic value (i.e., lowest  $LC_{50}$ ) for each chemical was selected as is done with the New Chemicals Program under TSCA and by Environment Canada
- Value was given for 349 out of 429 chemicals
  - Chemicals were excluded when it was predicted that the chemical may not be soluble enough to meet the predicted toxic effect (i.e.,  $LC_{50}$ )



## OncoLogic



- The OncoLogic program was run on each chemical that a structure was available for in the program
- The program assigns a baseline concern level from high to low for a chemical to have the potential to cause cancer
- The chemical analog structure activity method was used with some standard exposure scenarios selected



# OncoLogic Results



- OncoLogic
  - 146 chemicals were successfully run
    - » High = 0
    - » High-Moderate = 10
    - » Moderate = 24
    - » Low-Moderate = 34
    - » Marginal = 29
    - » Low = 49
- 32 chemicals had pre-existing cancer screening (i.e., IARC, NTP, etc.)



# Fate, Transport and Exposure Potential



- Divide Priority Chemicals into Chemical or Use Classes
  - e.g., Chemical classes – Fluorinated, Brominated, Chlorinated, Siloxanes,
  - e.g., Fragrances, rubber chemicals, solvents, chemical intermediates, flame retardants
- High release – plasticizers, hydraulic fluids, or solvents, fire fighting surfactants, fragrances
- Degraded during use – antioxidants, vulcanizing agents, UV stabilizers, polymer initiators
- Stable during use – auto wax, defoaming agents, soil repellent
- Analyze Physical Properties, Binding, Release
  - Collect from PHYSPROP or Estimate with EPI Suite
  - Determine if chemically bound (reactive flame retardants vs. non-reactive)
  - Integrate physical properties & form during chemical use to assess release (use past examples of chemical release of known contaminants)



# Pollution Prevention



- Need Variety of Approaches Dependent Upon the Use and Chemical Properties Required for the Application
  - Synthetic intermediate: develop different synthetic route
  - Solvent: examine physical properties (vapor pressure, solubility for application)
  - Surfactant: same surfactant properties but more biodegradable
  - Flame retardant: possibly use phosphate to quench flame rather than halogens
  - General: structurally redesign molecule to make less persistent (less fully halogenated, linear alkyl chains – more biodegradable) but keep essential properties for use



# The Emerging Challenge



Confirming SAR predictions with environmental measurements for priority PB&T chemicals

Instrument technology  
High resolution separations & mass measurement  
2D-GC-TOF, LC-MS/MS and LC-QTOF

Analytical Method development

- new or refinements of existing extraction and isolation procedures
- analytical standards and reference materials

Contaminant free reagents  
Clean room isolation

Bioanalytical Methods and directed bioassays

*"in silico"* technology – computational toxicology  
Improved QSARs to identify P,B&T parent and metabolites

e.g., metabolites via TIMES (tissue metabolism simulator), CATABOL, BIOWIN



# Challenges and Opportunities



- Our screening of 000s of substances e.g., DSL and TSCA Inventory has yielded some interesting probable P&B substances
- Uncertainties in this type of screening include:
  - Possibility of **false positives** and **false negatives**
  - Lack of information on uses and actual emissions of the chemicals
  - Need for information on degradation products
- Lack of information on **biological effects** of the selected chemicals
- Numerous opportunities for both environmental chemists and environmental toxicologists
  - Analytical and bioanalytical methodologies
  - Exposure and risk assessment
  - SAR development
  - Chemical fate modeling



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